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(54) Title: CELL-FREE SYNTHESIS AND ISOLATION OF NOVEL GENES AND POLYPEPTIDES

(57) Abstract

A method for the cell-free synthesis and isolation of novel genes and polypeptides is provided. Within one embodiment, an expression unit is constructed onto which semi-random nucleotide sequences are attached. The semi-random nucleotide sequences are fast transcribed to produce RNA, and then translated under conditions such that polysomes are produced. Polysomes which bind to a substance of interest are then isolated and disrupted; and the released mRNA is recovered. The mRNA is used to construct cDNA which is expressed to produce novel polypeptides.

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CELL-FREE SYNTHESIS AND ISOLATION OF NOVEL GENES AND POLYPEPTIDES

Technical Field

expressing semi-random DNA or RNA seguences, isolating novel genes from those sequences, and using those genes to create <u>vitro</u> and, more specifically, to methods of generating and synthesis and isolation of novel genes and polypeptides $\pm n$ The present invention generally relates to the novel polypeptides.

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BACKGROUND OF THE INVENTION

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microorganisms would have to be screened individually to locate unfeasible, unless the novel gene provides the organism with a 10^{13} possible permutations. If 10 of these permutations had a polypeptide string of 10 amino acids has $20^{10}\ \mathrm{or}$ approximately specific antigen), then a population of $10^{12}\,$ would have to be current state of the art, the 10^{12} independently transformed order to obtain the sequence(s) of interest. For example, a number of new sequences for a specific property is virtually The isolation of novel genes and polypeptides from screened for the expectation of finding one desirable novel gene. Through the use of conventional methods (expressing distinct growth or survival advantage. Indeed, under the screen a large, genetically diverse population of cells in novel genes via microorganisms), the screening of a large semi-random sequences is currently limited by the need to desirable characteristic (such as the ability to bind a that one desirable novel gene.

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novel gene products which are localized within cells, colonies derived from each transformed cell must be treated to break open the cells. Typically 1000-2000 bacterial colonies per standard petri dish are lysed (e.g., by chloroform) for the Within present screening procedures for detecting screening procedure. Thus, to examine $10^{12}\ \mathrm{transformed}$ 32

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logarithmically dividing cells may be necessary for producing organisms, 500,000 to 1 billion petri dishes would be necessary. In addition, 10,000 to 100,000 liters of the large numbers of transformable cells.

synthesis of a novel desirable polypeptide. However, even at a sorter over 60 years to screen 10^{12} cells. Thus, present day Alternatively, where a gene product is secreted and attached to the outside of a cell, it may be detected by its time-consuming, effectively prohibit the isolation of novel ability to bind a fluorescent compound or other marker. In flow rate of 5,000 cells per second, it would take a cell these cases, cell sorters may be used to screen for the screening methods which are both extremely costly and genes and polypeptide from semi-random sequences. 2

conformations for the method to have commercial applicability. for selectably obtaining polypeptides which specifically bind expressed in yeast at reasonably high levels and in "native" In addition to the methods briefly discussed above, Fields and Song (<u>Nature 340</u>:245-246, 1989) proposed a method gene. However, this system has serious limitations. First, to other polypeptides, using the domains of the yeast GAL4 both the known and novel binding polypeptides have to be polypeptide-nonpolypeptide interactions are excluded. only polypeptide-polypeptide binding may be selected; 20 5

diversity obviates the need for extremely large amounts of DNA, 4.5% of the control GAL4 activity. Fifth, Fields and Song used Fourth, it is not clear Whether random or semi-random sequences physical interactions were well-established and yet showed only very large sequences: 633 amino acids of the SNF1 protein and secondary structures that interact with each other. Sixth, using their method for semi-random sequences of even $10^{10}\,$ Third, glycosylated polypeptides or polypeptides that have special modifications may also be excluded by this method. 322 amino acids of the SNF4 protein, which have evolved can work, given that they used known polypeptides whose modifying enzymes, and competent yeast cells. 35 30 25

Contrary to previously disclosed methods, the present invention describes a method for cell-free screening of novel

associated with large numbers of transformed organisms as well as the limitations of the method disclosed by Fields and Song, methodology allows a substantial time and monetary saving in genes and polypeptides. This method avoids the problems and may be completed within a few weeks. Therefore, the the isolation of novel gene products.

SUMMARY OF THE INVENTION

methods for synthesizing, screening, and selecting high numbers polymerase binding sequence, a ribosome binding sequence, and a comprise the steps of (a) constructing an in vitro expression unit comprising a 5' untranslated region containing an RNA Briefly stated, the present invention relates to of novel genes and polypeptides. The methods generally

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expression unit and semi-random nucleotide sequences to produce transcribing or replicating the sequences associated with the conditions sufficient to maintain the polysomes; (e) binding semi-random nucleotide sequences to an expression unit; (c) translation initiation signal, the expression unit being RNA; (d) translating the RNA to produce polysomes under capable of producing mRNA; (b) attaching one or more 12

recovering and constructing cDNA from the released mRNA; and the polysomes to a substance of interest; (f) isolating the disrupting the isolated polysomes to release mRNA; (h) (1) expressing the gene to produce novel polypeptides. polysomes that bind to the substance of interest; (g) 20 25

through the various steps outlined above to further enrich for In one embodiment of the method described above, the significant (>10-3) fraction of the truncated population. In principle, the method may be repeated until the population of desirable sequences by amplifying the RNA or respective cDNA. desirable novel genes until desirable sequences represent a process may be repeated on mRNA that has been enriched for Subsequently, this amplified subset of genes may be cycled genes is nearly homogeneous.

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method for producing novel polypeptides is provided, comprising Within a second aspect of the present invention, a the steps of (a) constructing an in vitro expression unit

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polymerase binding sequence, a ribosome binding sequence, and a translation initiation signal, the expression unit being comprising a 5' untranslated region containing an RNA capable of producing mRNA; (b) attaching one or more

- polypeptides; (e) subdividing the RNA encoding the biologically transcribing sequences associated with the expression unit and semi-random nucleotide sequences to the expression unit; (c) semi-random nucleotide sequences to produce RNA; (d) translating the RNA to produce biologically active
- subdividing as set forth in steps (c)-(e) so that the gene of interest is isolated; (g) constructing cDNA from the isolated active polypeptides; (f) transcribing, translating, and gene; and (h) expressing the cDNA to produce novel polypeptides. 10
 - polymerase binding sequence, a ribosome binding sequence, and a method of producing novel polypeptides is provided comprising In yet another aspect of the present invention, a the steps of (a) constructing an in vitro expression unit comprising a 5' untranslated region containing an RNA 15 20
 - replicating the sequences associated with the expression unit and semi-random sequences to produce RNA; (d) translating the semi-random nucleotide sequence to the expression unit; (c) translation initiation signal, the expression unit being capable of producing mRNA; (b) attaching one or more
 - polypeptides; (f) translating and subdividing as set forth in steps (d)-(e) such that the gene of interest is isolated; (g) constructing cDNA from the isolated gene, and (h) expressing subdividing the RNA encoding the biologically active RNA to produce biologically active polypeptides; (e) the cDNA to produce novel polypeptides. 25

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translation initiation signal. The expression unit may further comprise a translation enhancer or "activator" sequences, a 3' The expression unit described above comprises an RNA polymerase binding sequence, a ribosome binding site, and a

DNA, by chemically synthesizing the DNA, or by polymerizing the The semi-random DNA sequences may be generated by mechanically, tail of a selected sequence and appropriate restriction sites. chemically, or enzymatically fragmenting naturally-occurring 35

organic polymer, active site of a protein molecule, metabolite, interest may be a surface antigen, receptor protein, toxin, DNA directly onto the expression unit. The substance of antibody, metal, hormone, or other compound.

These and other aspects will become evident upon reference to the following detailed description.

DESCRIPTION OF THE PREFERRED EMBODIMENT

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with commercially important properties, such as novel catalytic substances. Novel genes may be constructed which comprise open The present invention is directed to the isolation of advantageous to express the novel genes <u>in vitro</u>, as part of a virtually infinite diversity and may code for new polypeptides nucleotide sequences of chemically synthesized DNA. They may binding sites, and/or terminators. In some cases, it may be promoters, enhancers, initiation codons, plasmids, ribosomal be expressed in a wide variety of organisms using existing activities or the ability to bind selectively to specific novel genes and polypeptides. These novel genes may have reading frames from existing genes or from semi-random large-scale production process.

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multistep process for constructing and isolating novel genes As noted above, the present invention describes a preferred embodiment, the process comprises the following specific binding and/or biological activities. Within a and gene fragments which encode novel polypeptides with

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translation enhancer or "activator" sequences, and a 3' tail of An expression unit is constructed which contains expression unit may also contain convenient restriction sites, an RNA polymerase binding sequence (i.e., a promoter or an RNA-directed RNA polymerase initiation site), a ribosome binding site, and a translation initiation signal. The a selected sequence.

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fragmenting naturally-occurring DNA, RNA, or cDNA seguences, or by chemically synthesizing the nucleotides. The semi-random Semi-random DNA or RNA sequences are then generated by mechanically, chemically, or enzymatically

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The novel genes are then transcribed in vitro to polymerized directly onto the expression unit. A library of DNA or RNA sequences are then inserted into the expression $_{
m 10^{12}}$ or greater different sequences may then be created. unit. Alternatively, the semi-random sequences can be

an RNA-directed RNA polymerase sequence is included, then these produce a pool of RNA copies of the original DNA library. If

The RNA (mRNA) is translated in vitro to produce replicases may be used to amplify the RNA.

(RNA-ribosome-nascent polypeptide complexes) are used to keep polysomes. Conditions for maintaining the "polysomes" the desired polypeptide and mRNA together. 2

The polysomes are then allowed to bind to

hormones, and active sites of protein molecules, or to display proteins, toxins, organic polymers, antibodies, metabolites, substances of interest, such as surface antigens, receptor biological activity. 13

interest are substantially enriched by the removal of the unbound polysomes. Serial or flow-through washes under Polysomes binding to the substance(s) of

conditions which maintain the polysome complexes substantially increase the frequencies of the desired mRNAs, which remain attached to the substances of interest through the polysome structure. 20

The bound/active polysomes are then disrupted to release the mRNAs from the polysome complex. 25

The rare mRNAs are then recovered by making cDNA copies or by direct amplification of the RNA with RNA-directed RNA polymerases. The amplification of the CDNA with DNA

polymerase and/or reverse transcriptase reactions may allow The resulting cDNAs are then expressed to greater ease in recovering these low abundance messages. 30

produce polypeptides.

binding proteins above a background of nonspecific binding of preferable to further increase the frequency of specific In most instances, repetition of steps 3-8 is polysomes. 32

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The isolated, purified novel gene(s) produced by the methods described herein are capable of generating a variety of polypeptide(s) of interest using standard expression techniques, as positive proof that the gene codes for the desired product. In addition, DNA and/or polypeptide sequencing by conventional methods may be used to identify the composition of the novel polypeptide.

Once the polypeptide encoded by the novel gene has been isolated and identified, large-scale production of the novel polypeptide(s) may be accomplished by chemical synthesis (if the amino acid sequence is relatively short) or through recombinant DNA methods, using genetically engineered microorganisms. Alternatively, large-scale in vitro transcription and/or translation methods may be used to produce commercial quantities of the polypeptide.

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The DNA sequence coding for the selected polypeptide may also be incorporated into larger genes (i.e., such as into the hypervariable regions of antibody genes) to create hybrid proteins with the specific binding and/or biological activities of the originally isolated novel polypeptides, in addition to other binding and biological activities.

I. THE EXPRESSION UNIT

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The expression unit comprises a 5' untranslated 25 region and may additionally comprise a 3' region. The 5' untranslated region of the expression unit contains a promoter or RNA polymerase binding sequence, a ribosome binding sequence, and a translation initiation signal. The 5' untranslated region ("head") may also contain convenient

untranslated region (nead) may also concain convenient sequence(s). The 3' region may contain convenient restriction sites and a 3' tail of a selected sequence. The expression unit may be chemically synthesized by protocols well known to those skilled in the art. Alternatively, these elements may be incorporated into one or more plasmids, amplified in microorganisms, purified by standard procedures, and cut into appropriate fragments with restriction enzymes before assembly

into the expression unit.

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The 5' untranslated region contains a promoter or RNA polymerase binding sequence. High-efficiency promoters, such as those for the T7, T3, or SP6 RNA polymerase, are preferred in this invention for the following reasons. Such promoters

specific for their relative polymerases, and are highly active, allowing for greater than 50 rounds of transcription per DNA template. In addition, 77, 73, and SP6 polymerases are commercially available from many sources and are components of

10 well-characterized transcription kits. For the T7 promoter,
 the consensus sequence is TAATACGACTCACTATAGGGAGA (23 base
 pairs). Although this sequence is described in conjunction
 with a preferred embodiment of this invention, it will be
 evident that related DNA sequences may be used which will
15 function for T7 RNA bolymerase. and other sequences will be

15 function for T7 RNA polymerase, and other sequences will be appropriate for other RNA polymerases. Within certain embodiments, it may be desirable to utilize two promoters, such as both the T7 and SP6 promoters.

Positioned downstream of or within the promoter 20 region is a DNA sequence which codes for a ribosomal binding site. This ribosome binding site may be specific for prokaryotic ribosomal complexes (including ribosomal RNAs) if a prokaryotic translation procedure is used. However, a preferred embodiment of this invention uses a eukaryotic

25 sequence and an <u>in vitro</u> eukaryotic translation system, such as the rabbit reticulocyte system (Krawetz et al., <u>Can. J.</u> <u>Bicchem. Cell. Biol. £1:274-286</u>, 1983; Merrick, <u>Meth. Enzymol. 101</u>:38, 1983). A consensus translation initiation sequence, GCGCCCACCATGG, as well as other functionally related sequences

30 have been established for vertebrate mRNAs (Kozak, Nucleic Acids Res. 15:8125-8148, 1987). This sequence or related sequences may be used in the novel gene construction to direct protein synthesis in vitro. The ATG triplet in this initiation sequence is the translation initiation codon for methionine; in vitro protein synthesis is expected to begin at this point.

Between the promoter and translation initiation site, it may be desirable to place other known sequences, such as translation enhancer or "activator" sequences. For example,

untranslated region of alfalfa mosaic virus RNA 4 increases the Rueckert, <u>J. Virol. 37</u>:876-886, 1981), turnip mosaic virus, and severely reduce the expression of the SP6 RNAs (Jobling et al., virus "stimulated translation significantly" in SP6-generated Jobling et al. (<u>Nucleic Acids Res. 16</u>:4483-4498, 1988) showed that the untranslated "leader sequences" from tobacco mosaic interleukin mRNAs (Jobling and Gehrke, <u>Nature 325</u>:622-625, 1987). Black beetle virus (Nodavirus) RNA 2 (Friesen and efficiencies. In contrast, certain untranslated leaders brome mosaic virus coat protein mRNAs (Zagorski et al., mRNAs. They also reported that the 36-nucleotide 5' translational efficiency of barley amylase and human <u>Biochimie 65</u>:127-133, 1983) also translate at high

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NcoI site may also be used as a convenient cloning site for the For example, the sextuplet, CCATGG, is the recognition sequence Appropriate restriction sites may also be included in for the restriction endonuclease, NcoI. A NcoI "cutting site" site, and another promoter may be attached for expression $\underline{\mathbf{in}}$ the expression unit to assist in future genetic engineering. convenient splice point for subsequent genetic engineering. expression unit may be spliced from the novel gene at this polypeptide domains are brought together and expressed as <u>vivo</u> and large-scale production of the novel polypeptide. positioned downstream of the ribosomal binding site is a Hence, after purification of a desired novel gene, the construction of hybrid proteins, where two different single protein. ibid., 1988). 25 2 13

region of the gene (NotI is expected to cut totally random DNA coding region is dependent upon the nucleotide composition or cloning the novel gene into plasmids. The octameric sequence, GCGCCCGGC, is recognized by Noti nuclease and is particularly include in the 5' untranslated region a DNA sequence with at useful because it would rarely fall within the novel coding once every 65,536 base pairs). Other restriction sites may also be used; the expected frequency of cutting the novel least one restriction endonuclease site for subsequently In addition, it is most likely advantageous to

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the DNA source of the coding region. It should be noted that certain palindromic sequences may interfere with translation; however, some sequences may also enhance the rate of

The expression unit may also comprise a 3' region. translation.

engineering of the polypeptide coding region. For example, if thus, palindromes in the 3' region may slow down the movement of ribosomes during translation. This second property may be NotI "sticky ends" for further cloning. Second, palindromes desirable polypeptide coding sequence could be cut out with restriction sites would be convenient for any later genetic may cause secondary structures which impede translocation, It is desirable to construct known 3' regions (tails) with palindromic sequences for at least two reasons. First, 3' NotI sites were located in both the 5' and 3' regions, a 15 ដ

other polynucleotide stretch for later purification of the mRNA desirable for preventing ribosomes from "falling off" the mRNA translation step. The 3' region may also contain a poly-A or from other components in the in vitro translation reaction by and thereby enhancing the number of polysomes in the in vitro hybridization to a complementary homopolymeric sequence. 20

produced). One example is the 11 amino acid Substance P, which among the billions of novel polypeptides. The ID peptide would be useful for quantifying the amount of novel polypeptide and incorporated into the expression unit. Within one embodiment, semi-random amino acid sequences. The nonrandom component of acids (an identification or "ID" peptide) that is conserved nonrandom 5' untranslated region and/or with the 3' region. This nonrandom coding sequence specifies a string of amino can be attached as a fusion peptide to other polypeptides. for purification of the novel polypeptide (given that an In addition, other nonrandom sequences may be antibody against the ID peptide is available or can be the coding region is synthesized and produced with the the expressed polypeptides contain both nonrandom and 30 52

detecting and quantifying fusion proteins containing Substance Anti-Substance P antibodies are commercially available for 35

P. Another example is the eight amino acid marker peptide, "Flag" (Hopp et al., <u>Bio/Technology</u> 5:1204-1210, 1988). Amino-terminal ID peptides have at least two advantages over carboxy-terminal ID peptides. First, it is easier to make gene constructions which maintain the proper reading frame of the N-terminal ID, because long stretches of semi-random DNA or RNA will tend to end in all three reading frames for a C-terminal ID. Second, the N-terminal ID may be designed to function as a signal peptide in a transformed organism, allowing for the possible secretion of the novel polypeptide during large-scale production.

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Nevertheless, C-terminal ID polypeptides may also be used. One preferable C-terminal polypeptide is polyglycine, which is encoded by poly-dG and is read Gly-Gly-Gly, etc., regardless of the reading frame of the semi-random sequences. The polyglycine 3' end of the polypeptide may act as a noninterfering tether of the nascent peptide and allow the semi-random sequences greater access to bind molecules of interest. In addition, the poly-dG sequence may be used for priming second strand synthesis of the cDNA and may be useful for purification of the RNA or DNA with polyC or poly-dC. Other repetitive sequences, such as GGGCGGGC..., may be used to code for a recognizable peptide sequence which is expressed in all reading frames. A preferable form of the ID peptide is one which may be cleaved from the novel polypeptide by simple chemical or enzymatic means.

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In addition to the DNA expression unit, an RNA expression unit may be constructed for semi-random polypeptide synthesis. One possible advantage of the RNA expression unit is that the recovery of the polysomal mRNA does not have to go through an initial cDNA stage. Instead, the mRNA with the desired sequences may be amplified with an RNA-directed RNA polymerase, such as that of QB (Q Beta) replicase (Haruna and Spiegelman, Rroc. Nat. Acad. Sol. 54:579-587, 1965). This enzyme can make one billion copies of recombinant RNA in 30 minutes (Lizardi et al., Bio/Technology 6:1197-1202, 1988). One suitable cloning strategy for amplification of recombinant RNA is detailed in Lizardi et al. (1bid., 1988). For purposes

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of the present invention, other elements, such as restriction sites, enhancers, and ID sequences, may be added to the DNA plasmids which give rise to the QB RNA templates. Semi-random coding sequences may be inserted on these plasmids by standard

- DNA methodologies. When the QB replicase template is transcribed (for example, by T7 RNA polymerase), an RNA library capable of in vitro replication may be created which contains the semi-random gene sequences. Alternatively, a similar RNA expression unit may be constructed by chemically synthesizing the appropriate RNA molecules and assembling them via an RNA
 - 10 the appropriate RNA molecules and assembling them via an RNA ligase, such as the T4 RNA ligase (commercially available), which links together single-strand RNA and/or single-strand DNA.

15 II. SEMI-RANDOM NUCLEOTIDE SEQUENCES

Semi-random sequences of DNA or RNA are attached to the expression unit. Since the RNA expression units and semi-random sequences may be generated from a DNA template or constructed from chemically synthesized RNA or mRNA fragments in much the same manner as DNA expression units, the following description merely describes the process for semi-random DNA attachment to the expression unit. Those skilled in the art will readily be able to construct the RNA-equivalent of the expression units attached to semi-random polynucleotides.

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Semi-random DNA may be generated by at least three methods. First, naturally-occurring DNAs from virtually any living source may be mechanically, chemically, or enzymatically fragmented and attached to the 5' untranslated region with DNA ligase. Mixtures of fragments from different DNA sources may be used. The end result may be the selectable expression of an

- active "open reading frame" -- a portion (fragment) of a protein that has no "nonsense" (or "stop") codon, unless the activity resides in the extreme C-terminus of the molecule. In one embodiment of this invention, a gene coding for a known function may be fragmented; the resulting pieces are ligated to the 5' untranslated region and later screened for the
- the 5' untranslated region and later screened for the expression of activity in the polysome assay. By examining the smallest gene fragment which provides biological activity, an

peptides and hybrid therapeutic proteins and may be beneficial analysis may be useful for creating small biologically active for drug delivery, if smaller size assists the peptide in analysis of protein domains may be made. Gene fragment reaching the target site.

full-sized gene may be isolated through binding the polysome to A second method for generating semi-random DNA is to antibody, receptor protein, or other diagnostic molecule. The In another embodiment of the present invention, the cell-free expression of cDNA "fragments" as herein described from a cDNA library. By expressing cDNAs in vitro and using "fragmented" DNAs may be semi-randomly sized cDNA molecules may be orders of magnitude more sensitive than previously polysome selection, a very rare partial or perhaps even described methods in locating desirable CDNA clones.

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fragments and cDNA strategies described above, an active, open chemically synthesize the DNA. For example, relatively long sequences. "Open reading frame" implies that no stop codon exists and often indicates a sequence from within a protein synthesized with mixtures of nucleotides at each position. However, a statistical problem of nonsense codons becomes apparent with chemically synthesized DNA. For the gene reading frame is located from within existing protein DNA molecules of approximately 100 nucleotides may be 2 12

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Stop codons cause termination of translation and release of the strategies to reduce the frequencies of nonsense codons and to stretch coding for a string of 30 amino acids, the probability completely random DNA, with the equal likelihood of any of the nonsense codon is therefore 3/64 = 4.6875%. For a random DNA common amino acids at all positions may not necessarily have of at least one stop codon within that string is about 76%. TGA--represent three of the 64 possible DNA triplets. For synthesized DNA having enough diversity to code for all 20 nascent polypeptide from the ribosome complex. Therefore, four nucleotides in each position, the probability of a However, it should be noted that chemically open reading frames. The stop codons--TAA, TAG, and coding region. 35 30

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bypass the usual result of nonsense codons during protein translation are preferable, and discussed below.

particular to reduce the likelihood of nonsense codons. In the composition may be manipulated to favor certain codons and in synthesized with only C and T to theoretically avoid nonsense encoded. Lim and Sauer (Nature 339:31-36, 1989) have used an extreme case, the third position of each triplet codon may be codons. However, in this case not all 20 amino acids are More specifically, the A, T, C, and G base

position in synthesizing new regions of lambda repressor. This combination allows for any of all 20 amino acids at each codon positions and an equal mixture of C and G at the third codon and reduces the frequency of nonsense triplets to $1/32 \, = \,$ equal mixture of all four bases in the first two codon 2

In a preferred embodiment of this invention, unequal 3.125%. However, in a string of 30 amino acids the likelihood of at least one TAG stop codon is about 61%. អ

mixtures of the bases are used in all three codon positions to codon position the amount of A is reduced to half of the level of the other three bases. In the third codon position only ${\tt G}$ used, but only half that amount of T is used. In the second reduce the frequency of stop codons, while still allowing a high frequency of all 20 amino acids at all codons. In the first codon position equal molar amounts of C, A, and G are 20

result of this strategy is a greater than 79% probability that strategy. However, only tyrosine will be represented at less no stop codons will be present in a string of 30 amino acids. and C or G and T are used, and in equal molar amounts. The The proportions of the individual amino acids are slightly distorted in this case relative to a totally random DNA 25

than half of the expected frequencies compared to the random situation. 30

To further overcome the presence of nonsense codons

are underrepresented as the result of unequal mixtures of bases nonsense suppressing tRNAs be used in the <u>in vitro</u> translation eliminates all but the TAG stop triplet, and tyrosine codons when using chemically synthesized DNA, it is preferred that steps. In particular, since the strategy described above 32

codon have been well characterized -- as have many other nonsense tyrosyl-tRNAs will also be included in the translation step to at each codon position, a nonsense suppressor which recognizes tryptophane- or leucine-inserting suppressors of the UGA stop TAG (actually UAG in the mRNA) and inserts tyrosine into the manner that the tyrosyl-tRNA now "reads" UAG instead of the tyrosine-inserting nonsense suppressors may be generated by read the tyrosine codons. Nonsense suppressors can also be changing the anticodon region of a tyrosyl-tRNA in such a made for the other two nonsense codons. As an example, normal UAU and UAC tyrosine codons in mRNA. Normal growing polypeptide chain is most desirable. Such

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suppressors are known; and, therefore, the construction of such

molecules would be evident to those skilled in the art.

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suppressors. The nucleotide sequences of many nonsense

Nonsense suppressors of mammalian translation systems tyrosine-inserting UAG suppressor tRNA from yeast (Capecchi et suppressors has been reported as high as 70% in vitro (Pelham, Nature 272:469-471, 1978). Geller and Rich (Nature 283:41-46, nonsense codons in eukaryotic in vitro translation systems is Cell 25:497-506, 1981; Hudziak et al., Cell 31:137-146, 1982; 1985; Capone et al., EMBO J. 4:213-221, 1985; Diamond et al., al., Cell 6:269-277, 1975; Gesteland et al., Cell 7:381-390, are known (Burke and Mogg, Nucleic Acids Res. 13:1317-1326, possible with the use of suppressor tRNAs, including the different investigators have shown that the "reading" of Laski et al., EMBO J. 3:2445-2452, 1984). Additionally, 1976). Readthrough of the UAG stop codon by such yeast 1980) have successfully suppressed nonsense codons in 20 25

bacterial suppressor tRNAs and tRNA synthetase. Therefore, the premature release of polypeptides from the ribosomes during the particular, Pelham shows that a particular UAG codon in tobacco Furthermore, both Pelham (ibid., 1978) and Geller and Rich use of tRNA suppressors in the present invention to reduce (ibid., 1980) describe high levels of naturally-occurring reticulocyte systems with yeast suppressor tRNAs and with nonsense suppression in eukaryotic translation systems. translation step is well within the state of the art. 9 32

by "supraoptimal concentrations of Mg^{+2} ," or a reported 2.1 mM MgCl2. This level of magnesium ion or higher may therefore be mosaic virus may be "read" (suppressed) nearly 40% of the time used advantageously within the present invention to increase the readthrough of nonsense codons and to thereby reduce the problem of translation termination of longer semi-random nucleotide sequences.

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acid. For example, the elimination of G at position three in a codon prevents methionine and tryptophan from being included in used to strongly bias in favor of or against a particular amino the peptide. As another example, a nucleotide mixture which is different mixtures of bases at selected codon positions may be In generating the semi-random DNA by chemical means, biased toward a high-cysteine content may be desirable for producing short peptides with internal disulfide bonds for structural rigidity. Such rigid peptides may bind other molecules more tightly. 2 15

and extension with DNA polymerase and/or by including a poly-dx tail from which to prime with poly-dx'. Other methods, such as the use of terminal palindromes that create "hairpin loops" for self-priming, may be used for second strand synthesis. 100 μg molecules. If the semi-random synthesis strategy is used, the of double-stranded DNA of 100 nucleotides contains about 10^{15} nucleotide sequences may be accomplished by "random priming" Second-strand synthesis of these artificial 2

are expected to preserve the reading frame of the synthetic DNA merely provided for purposes of illustration; longer seguences coding potential exists within laboratory bench-scale amounts semi-random sequences of any given length. Shorter molecules of DNA. Such a synthetic DNA molecule of 100 nucleotides is chemically synthesized base is not 100%. Therefore, more may also be synthesized. In addition, shorter synthetic molecules may be generated and ligated together to make better than longer molecules, because each addition of 30 35

different polypeptide. Therefore, a very large diversity in

expectation is that each of these molecules codes for a

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nonsense codons may be avoided by the use of shorter artificial

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A third method for generating semi-random DNA is to DNA molecules. It RNA ligase or other means may be used to link together the short single-stranded DNAs.

preserves both the consensus vertebrate initiation site and the untranslated region. If no N-terminal ID sequence is used, the polymerization may occur immediately after the ATG initiation polymerize the molecules directly onto the 3' end of the 5' sequence or preferentially after the ATGG sequence--which Ncol site. The most commonly used enzyme for this

nonsense suppressing tRNAs may greatly assist in overcoming the to favor certain codons and reduce the frequencies of nonsense of dATP should reduce the frequencies of nonsense codons (TAA, instead of terminal transferase (A. Kornberg, <u>DNA Replication</u>, deoxynucleotide triphosphates. In particular, a lower amount TAG, and TGA). $E \cdot \underline{coll}$ DNA polymerase I is reported to carry Again, the A, T, C, and G base composition may be manipulated thymus), which is routinely used for generating homopolymeric codon by controlling the relative concentrations of the four enzymes or chemical methods may also polymerize DNA directly deoxynucleotide triphosphates, semi-random heteropolymers of onto the expression units. Second-strand synthesis is most out non-template (de novo) synthesis of DNA and may be used easily accomplished by random primer extension, but other W.H. Freeman & Co., San Francisco, Calif., 1980). Other problem of stop codons in this semi-random DNA sequence. polymerization is terminal transferase (usually from calf DNA may be synthesized on a DNA primer with a free 3'-OH. methods may provide the same result. Again, the use of regions for DNA cloning. However, by mixing different 25 20 5 2

III. TRANSCRIPTION OF THE NOVEL GENES

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If DNA expression units are used with the semi-random commercially available and extremely active. As an example, a seguences, mRNA may be easily created with RNA polymerase. As Approximately 50 mRNA copies may be synthesized routinely for DNA expression unit with a T7 promoter is treated with T7 RNA polymerase according to manufacturers' specifications. discussed above, T7, T3, and SP6 RNA polymerases are

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RNA copies of each different DNA molecule. An RNA library of 6 reflect the same level of diversity but now contain 50 or more $\mu_{
m g}$ may contain 50 copies of 10 12 different mRNAs that are each each DNA molecule in 30 minutes. The DNA may be degraded with RNase-free DNase. If the original DNA library had a sequence diversity of $10^{12}\,$ molecules, the resulting mRNA pool should acids. Since 6 μg is easily manageable in small test tubes, capable of expressing a semi-random polypeptide of 30 amino standard laboratory tools and vessels may be used.

mRNA may be generated during <u>in vitro</u> transcription (Hope and addition of diguanosine triphosphate "caps" (or analogs) for The 5' ends of mRNAs need to be modified with the efficient translation in eukaryotic systems. The 5' capped translation process (Krieg and Melton, N<u>ucleic Acids Res</u>. struhl, <u>cell 43</u>:177-188, 1985) and/or in the <u>in vitro</u> 9

example) is used during the RNA polymerization relative to GTP. 12:7057-7070, 1984). To cap messages during transcription, an (m7G(5')ppp(5')G, from Boehringer Mannheim Biochemicals, for available from Stratagene (California), which claims that An mRNA capping kit based on this method is commercially excess of diguanosine triphosphate or an analog thereof 20 5

If the expression unit is RNA-based, such as the QB 90%-95% of the resulting RNA is capped.

billion copies are easily attainable). However, the diversity replicase system, a few RNA copies may be generated with T7 or other promoter systems (see Lizardi et al., ibid, 1988) if the copies exist (or if the novel genes were assembled at the RNA virtually unlimited number of copies of the RNA library (one level), RNA-directed RNA polymerase is capable of making a novel gene constructions involve a DNA plasmid. Once RNA **5**2

the library may be self-sustaining at the RNA level without the of the library remains the same. With RNA phages, such as QB, necessity of going through a DNA intermediate. 30

IV. TRANSLATION OF THE RNA 35

known. For convenience, the rabbit reticulocyte or wheat germ Several in vitro translation methods are widely systems may be used with minor modifications. In vitro 19 translation kits are available commercially. For example, the

"Translation Kit, Reticulocyte, Type I" from Boehringer
Mannhelm Biochemicals has all components for 100 translation
reactions. Each reaction has been optimized for approximately
5 1 µg of mRNA in a 25 µl volume. One µg of mRNA is sufficient
to code for over 4 x 10¹² novel genes, as described above.
Therefore, it is possible to translate extremely high numbers
of novel genes in relatively small volumes. For example, 10¹³
805 ribosomes only weigh approximately 66 µg. Because of the
typected to saturate the mRNA, only a few ribosomes per message are
expected to saturate the mRNAs.

As described in the protocol for the representative translation kit noted above, GTP and m7G(5')ppp(5')G are required for the efficient translation of in vitxo transcribed 15 RNA. Even if mRNA capping has been previously performed during transcription, as described above, it may be advantageous to add the diguanosine triphosphate (or analog thereof) and guanylyltransferase (Krieg and Melton, ibid., 1984) to the translation reaction. In the absence of capping during translation of the two reagents are necessary for the efficient translation of the mRNA. In particular, when QB constructions are translated, diguanosine triphosphate (or analog thereof) and guanylyltransferase may be necessary for capping the RNA molecules during translation.

Other techniques may also be employed to optimize translation and especially ribosome attachment to the mRNAs. For instance, it may be desirable to add ribonuclease inhibitors, such as heparin. Eukaryotic systems, such as the wheat gexm and reticulocyte translation methods, may yield similar results to prokaryotic systems. The prokaryotic systems have the advantages of smaller ribosomes and more readily available nonsense suppressor tRNAs. In addition, in prokaryotic cells transcription and translation are often simultaneous reactions. In the absence of coupled transcription and translation in prokaryotes, mRNA stability is greatly reduced. Therefore, a prokaryotic in vitxo expression system may be used which combines transcription and

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As described above, a preferred embodiment of the present invention is the use of suppressor tRNAs (especially tyrosine-inserting suppressors), which may be produced through recombinant DNA technology and/or by the partial purification of these molecules from mutant cell lines. Radioactive amino acids, especially S15-methionine, may be useful for monitoring in vitro translation and for following low amounts of polysomes in subsequent steps.

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After about 30-60 minutes, protein synthesis begins
in the translation reactions. The precise time may be
determined for any given set of translation conditions by the
use of radioactive amino acids (such as S35-methionine) and
monitoring TCA precipitable counts, which is indicative of

- polypeptide synthesis. After the onset of protein synthesis, 15 cycloheximide at a final concentration of 1 µg/ml is added to prevent the movement of the ribosomes on the mRNAs (Lynch, Meth. Enzym. 152:248-253, 1987). This level of cycloheximide and a Mg⁺² concentration of 5 mM may be use to maintain the mRNA-80S ribosome-nascent polypeptide complexes (polysomes).
- inching the proposed instance polypering completes (polysomes).

 To other ribosome inhibitors may also be used since cycloheximide, for example, will not work on prokaryotic ribosomes. However, in the absence of GTP the polypeptide release from the ribosomes should not normally occur.

25 V. BINDING POLYSOMES TO SUBSTANCES OF INTEREST

The list of potential compounds to which the nascent peptide might bind is virtually unlimited. The coupling chemistries to link these compounds to columns, matrices, filters, beads, etc., will depend to a great degree upon the nature of the compound. In some cases, whole cells or cellular fractions may be used to find peptides which bind to cellular components, such as receptor proteins and other membrane-bound molecules.

For many proteins and nucleic acids, binding to 15 nitrocellulose or similar artificial surfaces is a property of the filters or fibres. In these cases, the substances of interest are "stuck" to the membranes by established protocols. Bovine serum albumin (BSA), gelatin, casein or nonfat milk, or

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VI. other proteinaceous material is then typically added in excess to bind up any "free" surface sites. For example, an antibody is first bound to nitrocellulose by placing a solution of the

dishes containing saline. After the washes, the disk is placed antibody on a nitrocellulose disk in a microtiter dish. After in a microtiter dish containing gelatin in solution. The disk washed by moving the nitrocellulose disk to fresh microtiter absorbing the antibody to the nitrocellulose, the disk is is then washed again with saline.

In this manner, polysomes which bind to the blocking protein or material (blocking protein) used in excess as described above. Before allowing the polysomes to bind substances of interest, it may be desirable to pre-absorb the polysome mix against BSA, gelatin, and in particular the proteinaceous

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polysomes binding to the substance of interest. For binding to specific antibodies (as in the case above), the pre-absorption step(s) may include another antibody, preferably of a similar subclass, but having different variable/hypervariable regions. By screening out polysomes which bind generally to antibodies invention may be useful for selecting anti-idiotypic binding pre-absorption step will lead to much greater specificity of activity (as seen for some anti-idiotypic antibodies) or be proteins. Such molecules may have biological or enzymatic but not to the variable/hypervariable region, the present nonspecifically to any protein are removed. This useful as vaccines. 22 20 15

or different pHs--may be used to locate polypeptides which bind inhibitors, such as heparin. In addition, specific incubation conditionally, depending on the environment. Incubation times parameters--such as low or high temperature, high or low salt, may be accomplished in the presence of ${\tt MgCl}_2$ (5 mM) and RNase will depend upon the concentration of the bound substance of The binding of polysomes to substances of interest interest and upon the nature of such substance.

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ISOLATION OF POLYSOMES WHICH BIND TO SUBSTANCE(S) OF

After allowing the polysomes to selectively bind to

conditionally-binding novel peptides may be isolated nonspecific binding of polysomes. If radiolabeled amino acids radioactive counts bound to the substance of interest. If the generally removed by washings. This wash should contain MgCl_2 amino acids are not labelled, washes should continue until at should continue until little detectable change is observed in are used in the translations, washes (serial or flow-through) and perhaps gelatin, BSA, or other proteins to help reduce least 10^{-6} dilution of the polysome solution is obtained. the substance(s) of interest, nonbinding polysomes are 2 വ

("stringent") condition(s) will be released into the solution and represent potential conditionally-binding factors against after these washes by shifting the polysomes into the desired concentration. Those peptides (and their attached ribosome different pH, high metal ion concentration, or low salt environment for nonbinding, such as higher temperature, mRNA complexes) which do not bind under the second 20 15

conditionally-binding peptides may be used to purify substances of interest. Alternatively, conditionally-binding peptides may serve as reagents in monitoring environmental changes. the substance of interest. Once immobilized,

VII. DISRUPTION OF THE ISOLATED POLYSOMES 25

The disrupted by the removal of ${\rm Mg}^{+2}$ (by dilution or via chelating agents) or through the destruction of proteins by a number of methods (proteases, chloroform, etc.). Although dilution is disruption of the polysomes as compared to other methods. bound polysomes are placed in a solution lacking ${
m Mg}^{+2}$ to the easiest method, it may not result in as thorough a The isolated (bound) polysomes may be easily liberate the mRNA; RNase inhibitors may be desirable. 30

Conditionally-binding polysomes, which were released under any of the desired environments, may be treated in a similar fashion to disrupt the polysomes and release their

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VIII. RECOVERING MESSENGER RNA AND CONSTRUCTING CDNA

Theoretically, if a single polysome binding to the substance of interest carries a mRNA, its rare mRNA is capable of being isolated (recovered) from the entire library of mRNAs. The mRNA may also be amplified by several techniques in order to facilitate isolation.

The use of the polymerase chain reaction (PCR) on a single copy of DNA and on rare mRNA is well documented. (For review, see H.A. Erlich (ed.), RCR Technology, Stockton Press, New York, N.Y., 1989; M.A. Innis et al. (eds.), RCR Protocols:

A Guide to Methods and Applications, Academic Press, San Diego, Calif., 1989; H.A. Erlich (ed.), Polymerase Chain Reaction:

Current Communications in Molecular Blology, Cold Spring Harbor Press, Cold Spring Harbor N.Y., 1989.) Briefly, the rare mRNA is first subjected to CDNA synthesis by standard means. Since the sequences of the 5' and 3' regions are known, specific

CDNA may then be amplified through the use of specified primers primers used for PCR may include sequences which restore the 5' recreating the expression unit in this manner, repeated rounds sequences which restore the promoter (e.g., the T7 polymerase simplified because each mRNA may be capable of replication to one billion copies or more, using the appropriate replicases. performed until virtually all of the selected genes code for primers may be used for cDNA synthesis. Second, the single binding peptides. For expression units based on RNA phages, such as QB, recovery and amplification of the rare mRNA is (even the same primers as those used in cDNA synthesis). and 3' regions of the original expression unit--that is, of transcription-translation-polysome selection may be recognition seguence) and 3' region are desirable. By 20 25 30

IX. EXPRESSION OF NOVEL GENES

Once the novel genes have been isolated and 15 sequenced, they or related sequences may be (1) cloned, (2) chemically reproduced, (3) mutated, and (4) expressed by protocols well known in the art. Large-scale production of the novel polypeptide may be accomplished through recombinant DNA

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transcription and translation produces a highly enriched peptide solution having the biological/enzymatic activity. At a frequency of 1 to 10^{-3} , the gene of interest may be readily isolated and cloned into appropriate expression systems, using methods currently available.

XII. CELL-FREE IDENTIFICATION OF NOVEL GENES AND PEPTIDES

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After a novel gene with putative binding or

biological activity has been isolated, it may be demonstrated
that the purified sequence codes for the activity of interest
by amplifying the DNA and/or RNA so that sufficient mRNA is
produced for larger-scale in vitz translation. The
translation products of this purified sequence should be nearly
homogeneous polypeptides having the assayable activity. The

15 gene and/or the polypeptide may be sequenced by existing methods to establish the composition of the novel polypeptide. Alternatively, the purified gene may be cloned into microorganisms for amplification and expression. Subsequently, biological/binding activities as well as sequence identity may

20 be established for the novel gene and polypeptide.

II. CREATING NOVEL HYBRID PROTEINS

After the nucleic acid sequence has been determined for the novel gene, this sequence may be incorporated into 25 larger genes to create hybrid proteins, which have the characteristics of the novel peptide and other desirable properties. One class of hybrid proteins which may be created by this technology is characterized by specific binding to cells and cytotoxic abilities. For example, a cell surface

or receptor-binding peptide may be joined to ricin or other toxins via DNA splicing methods. This type of hybrid protein may be used to selectively kill different cell types, including pathogens and tumor cells. The gene which encodes the hybrid protein may be completely synthesized or result from splicing the appropriate gene fragments together. This gene may be

expressed in a variety of expression systems.

A preferred embodiment of this invention is the replacement of variable and hypervariable regions of antibody

and antibody-like genes by novel gene sequences which code for binding activities against substances of interest. In this manner, a much greater range of diversity is possible against antigens of interest; and the screening process may be much more efficient and time-saving than the production methods for monoclonal antibodies against the same antigens. These "custom" hybrid antibody genes may be expressed in a number of organisms to produce active antibodies with new specificities or properties.

XIV. OTHER COMMERCIAL USES OF THE INVENTION

The application of the present invention in diagnostic tests parallels the use of monoclonal/polyclonal antibodies, and is more advantageous, primarily because the isolation of novel polypeptides as herein described may take considerably less time (one week versus a few months for antibodies). In addition, other advantages may be seen. The novel polypeptides may be considerably smaller molecules than novel polypeptides may be considerably subjection, and/or the antibodies. Therefore, synthesis, purification, and/or manufacturing of the novel peptides may be greatly simplified and cost-effective as compared to antibodies. The smaller size may also aid in stability, formulation, and in reaching the target molecules.

The novel polypeptides may be identifiable by (1)

fusing them to a biologically active peptide which has a
quantifiable activity (such as peroxidase or other enzymatic
activity), (2) synthesizing them with an ID peptide, described
above, to which existing antibodies are known to bind, (3)
above, to which existing them, (4) chemically adding markers,
radioactively labelling them, (4) chemically adding markers,

such as fluorescent dyes or metallic substances, or (5) any combination of the above. To increase specificity in the diagnostic use of the novel polypeptides, two or more different polypeptides may be used. In addition, novel polypeptides may be used as competitive binding elements in diagnostic tests be used as competitive binding to antigens or substrates.

35 which rely upon competitive binding to antigens or substrates.

Another advantage of novel polypeptides generated via

the present invention is that they may bind to many classes of

molecules which would not elicit a strong immune response,

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because some molecules are not complex enough or are too similar to an organism's resident compounds to trigger antibody formation. In addition, the use of novel polypeptides in diagnostic binding assays may have a much greater scope than the traditional antibody-based methods.

The novel polypeptides of the present invention may also be used therapeutically as originally isolated or as part of fusion proteins. For example, if a novel polypeptide were selected to bind a given toxin, it might also neutralize the toxin. If a new polypeptide is bound to a viral receptor site on a cell membrane or to the virus's attachment mechanism, infection of the cell may be diminished. As described earlier, fusion proteins carrying novel polypeptide recognition sequences in addition to a toxin may be used to selectively kill diseased or malignant cells. The binding of novel sequences to infected or malignant cells may trigger an immune response against the cell-peptide complex and, therefore, may be useful in the control of disease.

EXAMPLES

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The following examples are provided by way of illustration and not by way of limitation. Within the examples, standard reagents and buffers that are free from contaminating activities (whenever practical) are used. It is preferred to exercise care to avoid ribonucleases and PCR product contamination.

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EXAMPLE 1

SYNTHESIS OF A NOVEL GENE LIBRARY

gene library require careful planning by those skilled in the art. The 5' untranslated region of the expression unit contains an RNA polymerase site, a ribosome binding site, an initiation codon, and selected 5' untranslated sequences. The polymerase binding site used in this example is the T7 promoter sequence: TAATACGACTCACTATAGGGAGA (23-mer), which is placed at the 5' end of the expression unit.

A rabbit reticulocyte system is used for translation of the RNAs synthesized from the 77 promoter. Therefore, the ribosome binding site should include at least part of the consensus sequence for eukaryotic untranslated regions. In her review article, Kozak (ibid., 1987) suggests that very short untranslated regions (less than 10 nucleotides) do not initiate protein synthesis efficiently. A selected untranslated region of 36 nucleotides is used here. This untranslated region is derived from the naturally-occurring (36-base pair) upstream sequence of the adult rabbit hemoglobin (alpha-globin):

ACACTTCTGGTCCACTCACAAGGAACCACCATGG, where the underlined ATG represents the start of translation at a methionine initiation codon (Baralle, <u>Nature 267</u>:279-281, 1977). The rabbit alpha-globin untranslated sequence is chosen because (1) it is expected to be a favorable substrate in a rabbit reticulocyte system and (2) it contains the important "motifs" of Kozak's model mRNA.

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sequence, including the ATGG of the coding region, is therefore following ways for in vitro gene expression. First, the 5' A structure in the untranslated region of alpha-globin which is hypothesized to reduce the initiation of protein synthesis by This second change also creates a convenient GATC restriction 60% relative to the beta-globin mRNA (Baralle, ibid., 1977). (underlined above) is replaced by a G, which may aid in the capping of the mRNAs (Green et al., Cell 32:681-694, 1983). Second, the G (underlined in the alpha-globin sequence) is site in the 5' untranslated region. The resulting leader replaced with an A to help eliminate a putative secondary The alpha-globin sequence is modified in the the following: 20 25 30

 $\underline{g}_{\mathrm{CACTTCTG}}$ This leader sequence is placed immediately downstream from the T7 promoter.

The 3' region contains (1) a selected sequence for specific-primer-directed DNA synthesis, (2) a GGG-rich region which codes for a polyglycine tether that gives the nascent polypeptide spatial freedom to bind the substance of interest, and (3) convenient restriction sites whose resulting RNA

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secondary structure may impede the translocation of ribosomes off the mRNA. The polyglycine region comprises 20 codons for glycine; most of the glycine codons are adjacent GGG triplets, which code for glycines in all reading frames. However, some

5 of the glycine codons are GGT or GGA to keep the DNA strands in proper register. The restriction sites for Bam HI (GGATCC) and NotI (GGGCCGC) are chosen to be placed very near the 3' end of the gene; in the mRNA these sequences are expected to form hairpin loops. To prevent second-strand self-priming (of hairpin loops) by the NotI sequence, an addition of AAAA is made at the 3' end. The 3' region therefore has a general sequence of (GGG or GGT/A)₂₀ followed by GGATCGCGGCGCAAAA. A

The semi-random gene sequence is synthesized with 15 known 5' and 3' ends which undergo basepairing and ligation with the fully described 5' untranslated region and 3' region segments. To achieve this end, the semi-random gene is synthesized with a 5' CACCAIGG, which may basepair with the octamer CCAIGGTG on the complementary strand of the 5'

specific sequence for this region is given below.

outranslated region. The initiation (first) codon, <u>ATG</u>, is necessary for translation of the semi-random sequences. The subsequent G is the first position of the second codon and is constant to preserve the NcoI site at the front end of the gene. The rest of this second codon and the next 28 codons are

nonsense triplets. That is, in the first codon position, equal molar amounts of C, A, and G are used but only half that amount of T is used. In the second codon position, the amount of A is reduced to half of the level of the other three bases. In the third codon position, only G and C are used, and in equal molar and the codon position, only G and C are used, and in equal molar

amounts. After codon 30 is synthesized, GGTGGGGG is added. This sequence codes for two glycine residues and is used to ligate the semi-random sequences to the 3' region, which has a scomplementary CCCCCACC overhang on the opposite strand. The result of this synthesis is a sequence that codes for virtually all 30 amino acid polypeptides (beginning with methionine) and has a polyglycine tether. The probability of no stop codons in

partially purified yeast tyrosine-inserting UAG suppressor tRNA (Pelham, ibid., 1978) during the subsequent translation, over 90% of the semi-random sequences are expected to code for this string of triplets is approximately 80%. By using

The specific oligonucleotides to synthesize are full-length polypeptide.

listed below:

I. T7 Promoter & "Globin" Leader (for gene synthesis and PCR): S 'TAATACGACTCACTATAGGGAGAGCACTTCTGATCCAG

TCCGACTGAGAAGGAAC3'-OH

II. Anti-T7 Promoter & "Globin" Leader (for gene synthesis): 5 CCATGGTGGTTCCTTCTCAGTCGGACTGGATCAGAAGC

rcrccraracreagregrarra; -oH (5' kinased with T4 Polynucleotide 72

Kinase)

5.CACCATGG ... semi-random as described III. Semi-Random Gene (for gene synthesis):

GGTGGGGG3'-OH (5' kinased with T4 Polynucleotide Kinase) 20

IV. Poly-Glycine $\mathfrak k$ 3' Restriction Sites (for gene synthesis): 5 'TGGGGGTGGTGGGGGGGGGGGGAGGAGGGGGGG

GGGGAGGGGGAGGTGGTGGATCCGGGCGGCAAAA3'-OH (5' kinased with T4

Polynucleotide Kinase) 22 v. Anti-Poly-Glycine & 3' Sites (for gene synthesis): 5'ITITGCGGCGGGGATCCACCACCTCCCCCTCCCCCCCCC

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VI. Anti-Poly-Glycine & 3' Sites (for cDNA synthesis and PCR): 5 · TTTTGCGGCCGCGGATCCACCACCTCCC3 · -OH

longer, are allowed to cool slowly to room temperature, and are Seguences I and II are mixed in equimolar amounts in complementary sequences (which comprise the 5' untranslated region) are allowed to anneal at 50.-60.C for one hour or standard TE Buffer and heated at 65°C for 5-10 min. The 35

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treated to form the double-stranded 3' region. These duplexes each have an eight-base, single-stranded overhanging sequence thereafter stored at 0.-4.C. Sequences IV and V are likewise Equimolar amounts of I/II duplex, IV/V duplex, and which is complementary to the known ends of Sequence III.

ligation product, which is approximately 200 base pairs (233 bp Complete double-stranded synthesis of novel genes is if completely double-stranded, which it is not). The "200 bp" overnight at 13.-15.C in Ligase Buffer. The ligation mix is DNA band is gel purified with NA45 paper (S&S) or by any of then run on a 1.5% agarose gel to separate out the desired approximately $10^{13}\,\mathrm{DNA}$ molecules) or more is desirable. semi-random Sequence III are ligated with T4 DNA ligase several protocols. A total of 2.5 μg (representing ព

the "second-strand" synthesis of the semi-random sequences. T4 methods. The double-stranded 3' region provides a primer for Sequence II, thereby filling the nick in the second strand. The DNA library is phenol/chloroform extracted and ethanol accomplished with DNA Polymerase I, Klenow, using standard DNA ligase is used to join the newly synthesized DNA to 12

precipitated.

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10 μg of completely double-stranded DNA molecules has transcribed with T7 RNA Polymerase to yield translatable mRNAS. However, with each transcription, the DNA library is consumed, a sequence diversity of 4 imes 10¹³. This library may then be

unless DNA copies are made. To replicate the DNA library, 100 Technology, pp. 18-19 (Erlich, ibid., 1989), each 200- μ l PCR ng aliquots are each distributed to $500-\mu l$ tubes for PCR amplification in 200- μ l reactions. According to \overline{PCR} 25

pooled. The pooled sample contains on the average 50 copies of each translation with T7 RNA Polymerase. If the library is to each semi-random sequence and therefore may be used repeatedly (50 times, for example) without a large loss of diversity for 50-fold duplication of DNA in each aliquot. The aliquots are reaction yields about 5.2 µg of DNA--or an approximately 33 30

be replicated with PCR, then the Klenow filling and ligation steps, described above, may be unnecessary, since the Taq polymerase is capable of filling in the gap and

Examples of oligonucleotide primers for PCR amplification of the DNA library are listed above in sequences I and VI. Generally, oligonucleotides of 25-30 bases are used for PCR amplification; however, longer primers may be used. It is important that the primers do not share significant

10 homologies or complementary 3' ends. Sequences I and VI have noncomplementary ends and no obvious regions of extensive homology.

In addition, after translation of these novel gene

sequences, the resulting mRNAs lack T7 promoter sequences.

Sequence VI is used as the primer for first-strand cDNA synthesis. Sequence I is used as the primer for second-strand synthesis and restores the T7 promoter to the cDNA. In this way, later rounds of translation are possible on the selected novel gene sequences. PCR amplification may be necessary if the resulting cDNAs are relatively rare.

EXAMPLE 2

TRANSCRIPTION OF NOVEL GENES

The DNA library (or a representative aliquot of those sequences) described in Example One is transcribed with T7 RNA polymerase. 2.5 µg of this DNA codes for nearly 10¹³ different polypeptides. The DNA is capped during transcription with Stratagene's mCAP" Kit, according to the manufacturer's specifications. Approximately 5-10 µg of mRNA is expected.

30 Generally, with T7 RNA polymerase, nearly 10 times this level of RNA is synthesized; however, the conditions for the capping reaction limit mRNA production in this case. The DNA is removed with DNase I, provided in the kit. The capped mRNA is phenol/chloroform extracted and precipitated with ethanol. The SRNA is resuspended in 10 µl of TE and stored at 0--4°C.

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EXAMPLE 3

TRANSLATION OF NOVEL GENES

The capped mRNA is translated with Boehringer Mannheim Biochemical's rabbit reticulocyte kit, with all 20 amino acids at 312.5 μ mol/l each. Capped mRNA from Example 2 is added to each reaction at 0.5 μ g per reaction and is treated according to the manufacturer's protocol. After around 60 minutes at 30.°C, cycloheximide is added to a final concentration of 1 μ g/ml. MgCl₂ is adjusted to 5 mM, and

10 heparin is added to 0.2 mg/ml. The reactions are pooled and submitted to a discontinuous sucrose gradient, according to Lynch (ibid., 1987). The polysomes may be frozen at -70°C or used directly.

EXAMPLE 4

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IMMOBILIZATION OF ANTIBODIES

AS THE SUBSTANCE OF INTEREST

Antibodies may be used to select for novel binding peptides. Peptides which bind to the hypervariable/variable of regions of the antibodies ("anti-id peptides") may behave like the original epitopes which were used as immunogens. Because the novel anti-id peptides may mimic the original epitopes, these peptides may be useful as vaccines and/or may demonstrate biological activities, in much the same way that anti-id

25 antibodies have been shown to have biological (sometimes catalytic) activities.

Examples of useful antibodies are anti-fibronectin, anti-nerve growth factor, anti-CD4, and anti-tumor necrosis factor, which are all available from Boehringer Mannheim

Blochemicals. In general, antibodies to receptor molecules, growth factors, surface antigens, and biologically active peptides, as wall as neutralizing antibodies to toxins and diseases, are good candidates for which to isolate anti-id binding peptides that may have agonist or antagonist properties

The antibodies are affixed to Immobilon PVDF (polyvinylidene difluoride) membrane from Millipore Corporation, according to Pluskal et al. (Biolechiques

or serve as vaccines.

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clone 3E3, Boehringer Mannheim Biochemicals) is absorbed onto a

4:272-283, 1986). For example, anti-fibronectin antibody (from

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EXAMPLE 6

RECOVERING NOVEL GENES WHICH CODE FOR

ANTI-ID PEPTIDES FROM POLYSOMES

The PVDF is removed, placed in a fresh tube of 0.1 mM EDTA, and 100 μl of 0.1 mM EDTA and is gently shaken at room temperature antibody-bound polysomes, is transferred to a tube containing for 5-10 minutes to disrupt the polysomes and liberate mRNA. The PVDF membrane, which holds the washed

transcribed; and the resulting cDNA is amplified, according to Instead of using random hexamer for priming the cDNA synthesis PCR Technology (ibid., 1989), p. 91, with slight modification. a sequence complementary with the known 1' region (such as stored at 0.-4.c overnight or longer (as a back-up). The released mRNA from the first EDTA treatment is reverse 10

amplified as described in PCR Technology, using Sequence I or a similar DNA upstream primer. After PCR amplification, the five 20-µl reaction). After the reverse transcriptase reaction, the appropriate relative amounts of the other reagents (instead of aliquots are pooled, phenol/chloroform extracted, and ethanol Sequence VI listed earlier as the downstream primer) is used transcriptase step is done in 100 μl of PCR buffer with the mixture is split into 20 μl aliquots; and each aliquot is for both cDNA synthesis and PCR reactions. The reverse 20 12

sites of the PVDF. The membrane is then washed twice with 0.1%

temperature, so that the gelatin is absorbed into unoccupied

gelatin in saline buffer. A similar treatment is done with 10

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 μg anti-keratin antibody (from clone AE1, Boehringer Mannheim

Biochemicals), which is the control IgG_1 as described below.

in saline buffer is absorbed onto the PVDF square by incubating

at room temperature for at least two hours. The PVDF is then

washed with the Saline Buffer twice. The membrane is next

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incubated with a "blocking solution," containing 5% (w/v) gelatin in saline buffer for at least two hours at room

 $\mu_{
m g/cm}^2$ of IgG. For convenience, 1 $\mu_{
m g}$ of anti-fibronectin IgG $_{
m L}$

anti-id peptides(s); Immobilon" pvDF is reported to bind 172

needed is dependent upon the binding parameters of the desired

100% methanol and washed twice with 0.9% (w/v) NaCl in 10 mM

0.5 cm \times 0.5 cm square of PVDF, that has been "wetted" with

Tris buffer pH 7.4 (Saline Buffer). The amount of antibody

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precipitated. This cDNA is then resuspended in TE and stored at 0.-4.C. 25

expression methods are practical. In addition, by dilution to previously described. In this case, the desired sequences are low Poisson Distribution of genes, a single novel gene(s) may repetition of this cycle, which is greatly aided through the use of programmable workstations, desirable novel genes are greatly amplified compared to the original DNA library. be isolated, amplified, transcribed, and translated to concentrated to a level where conventional cloning and polymerase and translated in a reticulocyte system, as The selected DNA is transcribed with T7 RNA 35 30

under the same conditions to allow specific polysome binding to

antibody. The anti-fibronectin PVDF square is removed and washed three times by transferring it serially to fresh PS

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the variable/hypervariable region of the anti-fibronectin

gelatin and \log_1 . The anti-keratin PVDF square is removed with

jewelers' forceps and is replaced with the anti-fibronectin

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PVDF square. The mixture is incubated for four more hours

pre-absorption step is done at 0.-4.C with gentle agitation for

 μg anti-keratin IgG, described in Example 4. This

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four hours to select out nonspecific binding of polysomes to

heparin, and 1 $\mu g/ml$ cycloheximide) and a PVDF square with 10

incubated in 1-ml reactions, each containing PS Buffer (0.9%

Polysomes with nascent semi-random peptides are

POLYSOME BINDING TO ANTIBODIES

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NaCl, 10 mM Tris pH 7.4, 1% gelatin, 15 mM MgCl2, 0.2 mg/ml

demonstrate specific binding capability of the gene product(s).

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Once binding has been demonstrated, the isolated gene(s) and polypeptide(s) may be sequenced for identification.

After the sequence of the novel binding peptide is known, many methods exist for the manipulation and large-scale synthesis of the peptide, as described herein.

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EXAMPLE 7 COMPETITION ASSAY FOR BINDING PEPTIDES

After novel genes which code for binding peptides are selected, the amplified pools of recovered cDNA are assayed for the presence of the genes. Where ID sequences have been intentionally included to be coexpressed with the semi-random DNA sequences, ELISAs or other immunological assays for the known part of the peptide are used to detect the binding of the novel portion of the peptide to the substance of interest. However, when no ID sequence is present and/or a confirmation of binding specificity is desirable, competition assays for the peptides are carried out. Competition assays, including competition ELISA tests, are used to monitor for the presence of binding sequences within the various cDNA pools generated by the present invention.

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One example is the screening of the CDNA pools for genes which encode peptides which bind anti-Pseudomonas exotoxin (anti-PE) antibody. After two rounds of selection for polysome binding to the anti-PE antibody, different aliquots of the resultant CDNA pool were each transcribed in a $200-\mu l$ reaction with T7 RNA polymerase (30 units) under standard conditions, starting with approximately 200 ng of DNA. The mRNA products were phenol/chloroform/isoamyl alcohol extracted and precipitated with sodium acetate and ethanol at -20° . The precipitates were each centrifuged and resuspended in $16~\mu l$ of distilled water which had been treated with distilled water which had been treated with

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The resuspended mRNAs were heated to 65° for five minutes and then placed on ice. The RNAs were translated with a wheat germ kit (Boerhinger Mannheim Biochemicals) according to the manufacturer's recommendations. Each RNA sample was

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expressed in a 50 μ l translation reaction with 25 microcuries of 355-methionine and 0.5 μ l of RNase inhibitor. The reactions were run for 15-60 minutes at 30°. At the end of the

- translation, the samples were each equally divided: one half
 s was used to bind the substance of interest without competing
 substrate, while the other half was used to bind the substance
 of interest in the presence of excess competing substrate. In
 this case, the substance of interest was anti-PE antibody. The
 competing substrate was a 14-amino acid peptide (PE peptide),
 - 10 which is derived from the toxin protein sequence and known to bind the antibody. The PE peptide sequence is Val-Glu-Arg-Leu-Gun-Ala-His-Arg-Gln-Leu-Glu-Glu-Arg. See Wozniak, et al., Proc. Natl. Acad. Sci., 85: 8880-8884 (1988).
- The competition assays were done over ice in 96-well microtiter dishes with flat bottoms. Immobilon PVDF disks were made with a standard 1/4 inch holepunch and placed in wells labelled "A". 50 μ l of methanol were added to the disks in "A" to wet and sterilize the membranes. The disks were transferred with forceps to wells "B" which contained 200 μ l of Saline Buffer plus 10 mM MgCl₂ (TSM buffer). The disks were further washed by moving them to wells "C" which also contained 200 μ l
 - O Buffer plus 10 mM MgCl₂ (TSM buffer). The disks were further washed by moving them to wells "C" which also contained 200 μl of TSM. They were then transferred to wells "D" which contained 25 μl TSM plus 3 μl of anti-PE antibody (4.6 μg/μl). The antibody was absorbed to the disks for three hours on ice
- 25 with gentle rotation (50-100 RPM on a platform shaker). Afterwards, 75 μ l of 2% nuclease-free BSA was added to "D" and absorbed for 1 hr. at 100 RPM.
- The disks were washed twice in 200 μ l of TSM plus 0.1% BSA (in wells "E" and "P") for 30 minutes in each well and 30 were then ready for peptide binding. In wells "G" 26 μ l of TSM plus 0.1% BSA was mixed with 25 μ l of each translation reaction described above-half of the 50 μ l wheat germ system. Into one-half of each of the "G" wells, 1 μ l of PE peptide (1 mg/ml in TSM) was added to competitively inhibit the binding of novel sadioactively-labelled peptides to the antibody; these wells were labelled "+ Peptide." Into the control "G" wells, 1 μ l of

ISM was added and the wells labeled: "No Peptide." The disks were added to the appropriate "G" wells and incubated to three

antibody.

each disk was measured in a liquid scintillation counter with a obtained from the binding of polysomes to the anti-PE antibody: minute incubation for each wash. The bound radioactivity for results of competition assays on different aliquots of cDNAs After the binding reaction each disk was serially washed eight times in 200 μl of fresh TSM at 0°, with a l0 1 ml cocktail of Ecoscint. The following table lists the

CPM 35S-MET 5792 6303 8337 7693 6969 6163 WP1, No Peptide WE2, No Peptide WE1, No Peptide WP1 + Peptide WE2 + Peptide WE1 + Peptide SAMPLE 20 20 15

5845 6398

WP2, No Peptide

WP2 + Peptide

products of the selected cDNA pools, compared to the No Peptide amount of binding of the radioactively-labelled translation In each case the competing PE peptide reduced the plasmids, such as pUC18, pUC19, Bluescript, and many other sequences which code for binding peptides to the anti-PE sequences is then done by cloning individual genes into controls. These results indicate the presence of gene antibody. Isolation and characterization of these DNA available vectors. 30 25

modifications may be made vithout deviations from the spirit described herein for the purposes of illustration, various From the foregoing it will be appreciated that, although specific embodiments of the invention have been 35

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and scope of the invention. Accordingly, the invention is not to be limited except as by the following claims.

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WHAT IS CLAIMED IS:

- A method for producing novel polypeptides, comprising:
- and a polymerase binding sequence, a ribosome binding sequence, translation initiation signal, said expression unit being constructing an in vitro expression unit comprising a 5' untranslated region containing an RNA capable of producing mRNA; (a)
- (b) attaching one or more semi-random nucleotide sequences to said expression unit;

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- associated with the expression unit and semi-random nucleotide transcribing or replicating the sequences sequences to produce RNA; <u>0</u>
- (d) translating said RNA to produce polysomes under conditions sufficient to maintain said polysomes; 15
 - (e) binding said polysomes to a substance of
- interest;
- isolating said polysomes that bind to said (£)
 - substance of interest;

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mRNA;

- disrupting said isolated polysomes to release <u>6</u>
- recovering said mRNA; E
- constructing cDNA from said recovered mRNA; and $\overline{\Xi}$
 - expressing said cDNA to produce novel Ð

polypeptides.

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The method of claim 1 wherein, subsequent to the step of recovering mRNA and constructing cDNA, amplifying said cDNA by polymerase chain reaction.

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- The method of claim 1 wherein said semi-random nucleotide sequence comprises deoxyribonucleic acid.
- The method of claim 1 wherein said semi-random nucleotide sequence comprises ribonucleic acid. 35

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- The method of claim 1 wherein said expression unit includes at least one RNA-directed RNA polymerase recognition sequence.
- The method of claim 5 wherein said RNA-directed RNA polymerase is Q-Beta replicase. ė.
- The method of claim 1 wherein, subsequent to the step of recovering, amplifying the mRNA. 7.
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- amplifying comprises synthesizing duplicate sequences with an The method of claim 7 wherein the step of RNA-dependent RNA polymerase.
- The method of claim 8 wherein the RNA-dependent RNA polymerase is Q-Beta replicase. ٠ م 15
- isolating comprises removing polysomes that do not bind to said substance of interest by serial dilution or flow-through wash 10. The method of claim 1 wherein the step of

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- step of isolating said polysomes, said polysomes are exposed to 11. The method of claim 1 wherein, subsequent to the
 - selected stringency conditions such that said polysomes are released from said substance of interest. 25
- exposing said polysomes comprises raising the temperature, 12. The method of claim 11 wherein the step of lowering the salt concentration, or raising the metal ion concentration of said polysomes. 30
- 13. A method for producing novel polypeptides, comprising:
- polymerase binding sequence, a ribosome binding sequence, and constructing an in vitro expression unit comprising a 5' untranslated region containing an RNA 35

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translation initiation signal, said expression unit being

capable of producing mRNA;

(b) attaching one or more semi-random nucleotide

expression unit and semi-random nucleotide sequences to produce transcribing the sequences associated with the sequences to the expression unit;

RNA;

(d) translating said RNA to produce biologically

active polypeptides;

(e) subdividing the RNA encoding said biologically

active polypeptides;

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set forth in steps (c)-(e), such that the gene of interest is (f) transcribing, translating, and subdividing as

isolated;

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(g) constructing cDNA from said isolated gene; and expressing said cDNA to produce novel

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polypeptides.

14. A method for producing novel polypeptides,

comprising: 20

polymerase binding sequence, a ribosome binding sequence, and a translation initiation signal, said expression unit being constructing an in vitro expression unit comprising a 5' untranslated region containing an RNA

(b) attaching one or more semi-random nucleotide capable of producing mRNA;

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sequences to the expression unit;

expression unit and semi-random nucleotide sequences to produce replicating the sequences associated with the છ

RNA; 30

translating said RNA to produce biologically

(e) subdividing the RNA encoding said biologically active polypeptides;

translating and subdividing as set forth in active polypeptides;

steps (d)-(e) such that the gene of interest is isolated;

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constructing cDNA from said isolated gene; and

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expressing said cDNA to produce novel (F)

polypeptides.

The method of claim 14 wherein, subsequent to 15.

sequences associated with the biologically active polypeptides the step of subdividing the RNA, amplifying the novel gene with polymerase chain reaction or with an RNA-directed RNA ß

polymerase.

The polypeptide produced by the method of claim ដ

1, 13 or 14.

17. The method of claim 1, 13 or 14 wherein said ribosome binding site comprises eukaryotic, prokaryotic, or

viral ribosome binding sequences.

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translation initiation sequence, GCCGCCACCATGG, or functionally ribosome binding sequence comprises the vertebrate consensus 18. The method of claim 1, 13 or 14 wherein said

related sequences.

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selected amino-terminal ID peptide, said sequence positioned at expression unit further comprises a sequence.which codes for a 19. The method of claim 1, 13 or 14 wherein the

the initiation codon.

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consisting of sequences enhancing the amplification, cloning, replication, purification, and isolation of the novel genes. expression unit further comprises a 3' region of a selected The method of claim 1, 13 or 14 wherein said sequence, said selected sequence selected from the group 20.

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21. The method of claim 20 wherein said 3' region includes palindromic sequences which are adapted to impede

ribosome translocation. 35 22. The method of claim 20 wherein said 3' region includes a C-terminal ID sequence.

- 23. The method of claim 22 wherein said C-terminal ID sequence comprises a repetitive sequence.
- 24. The method of claim 22 wherein said C-terminal ID sequence codes for a peptide capable of binding to antibodies.
- expression unit further comprises restriction sites adapted to The method of claim 1, 13 or 14 wherein said allow expression of the novel gene in vivo. ខ្ព
- 26. The method of claim 25 wherein at least one of said restriction sites comprises the sequence CCATGG, said sequence positioned at the start of translation.

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expression unit includes the promoter seguences for T7, T3, or 27. The method of claim 1, 13 or 14 wherein said SP6 polymerase.

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- semi-random nucleotide sequences are generated by mechanically, chemically, or enzymatically fragmenting naturally-occurring 28. The method of claim 1, 13 or 14 wherein the DNA or CDNA.
- semi-random nucleotide sequences are generated by chemically The method of claim 1, 13 or 14 wherein the synthesizing nucleotides to form gene sequences.

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synthetically synthesizing said nucleotides comprises the steps of (1) utilizing substantially equal molar amounts of C, A, and in the first codon positions; (2) utilizing substantially equal G, and only half of said substantially equal molar amount of T positions; and (3) utilizing substantially equal molar amounts substantially equal molar amount of A in the second codon 30. The method of claim 29 wherein the step of molar amounts of C, T, and G, and only half of said of C and G or T and G in the third codon positions. 30 35

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- The method of claim 1, 13 or 14 wherein the step directly onto the 3' end of the 5' untranslated region of the of attaching further comprises polymerizing said nucleotides expression unit. 31. ß
- The method of claim 1 or 13 wherein the step of transcribing comprises transcribing said sequence in the presence of diguanosine triphosphate or analogs thereof. 32.

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- The method of claim 1, 13 or 14 wherein the step presence of diguanosine triphosphate or analogs thereof and of translating comprises translating said sequences in the guanylyltransferase. 33.
- The method of claim 1, 13 or 14 wherein the step of translating is conducted in the presence of nonsense-suppressing tRNAs. 34.

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- nonsense-suppressing tRNA is a tyrosine-inserting, The method of claim 34 wherein the nonsense-suppressing tRNA. 35. 20
- surface antigens, receptor proteins, toxins, organic polymers, substance of interest is selected from a group consisting of 36. The method of claim 1, 13 or 14 wherein said metabolites, active sites of protein molecules, hormones, antibodies, and pollutants. 25
- substance of interest is the variable/hypervariable region of 37. The method of claim 1, 13 or 14 wherein said an antibody. 30
- The method of claim 1, 13 or 14 wherein said substance of interest is a receptor protein. 38. 35
- 39. The method of claim 38 wherein said receptor protein is a growth factor receptor protein.

factor receptor protein is selected from the group consisting The method of claim 39 wherein said growth of insulin and epidermal growth factor. 40.

substance of interest is selected from the group consisting of 41. The method of claim 1, 13 or 14 wherein said viral surface antigen, viral receptor protein and CD4.

42. The method of claim 1, 13 or 14 wherein the step of expressing cDNA comprises chemically synthesizing the amino acid sequence based on the nucleotide sequence of said cDNA. 2

43. The method of claim 1, 13 or 14 wherein the step of expressing cDNA comprises cloning the nucleotide sequence into an expression vector for synthesis in genetically engineered microorganisms. 15

44. The method of claim 1, 13 or 14 wherein the step of expressing cDNA comprises in vitro transcription and/or translation of the nucleotide sequence. 20

encoding a polypeptide substantially homologous to that encoded sequence being substantially identical to the binding region of 45. The method of claim 1, 13 or 14 wherein the step of expressing cDNA comprises synthesizing a nucleotide seguence by said cDNA, the polypeptide encoded by said nucleotide said polysomes that bind to the substance of interest. 25

antibodies, enzymes, biologically active peptides, and peptides cDNA is joined to other selected nucleotide sequences selected 46. The method of claim 1, 13 or 14 wherein said from the group consisting of sequences encoding toxins, capable of binding to antibodies. 30

47. A method for isolating a nucleotide sequence which encodes a polypeptide of interest, comprising

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comprises a 5' untranslated region containing an RNA polymerase binding sequence, a ribosome binding sequence, a translation initiation signal, and one or more semi-random nucleotide transcribing an in vitro expression unit which sequences to produce a mRNA library;

and isolating mRNA from polysomes that specifically bind to the maintain polysomes having polypeptide chains attached thereto; contacting the polysomes to a substance of interest translating the mRNA library under conditions which

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substance of interest. ដ

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INTERNATIONAL SEARCH REPORT

1,3,4,10-14. 16,17,19,20, 22,24-26, 28-47 1,3,4,10-14, 16,17,19,80, 22,24-26, 28-47 DATABASES: Dialog Service Online (Files, 1967-1990; Automated Patent System 435/69.1; 435/69.7; 435/91; 435/172.1; 435/172.3; 536/27; :.. 530/350 Internetional Application No PCT/US90/05682 See Attachment BIOLOGY, Vol. 61, issued 1983, Krawetz et al., "In vitro Translation of Elastin mRNA and Processing of the Translated Products and the Signal Sequence Of Elastin a." pages 274-286, see entire document. III. DOCUMENTS CONSIDERED TO BE RELEVANT!!
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"X" deciment of particular orderance; the claimed immediate interests to receive the condered to increase an investment of particular independent; the claimed immediate independent of the claimed immediate properties of the condered to increase the owner the endered the condered out increase the sense of deciment in combined with one of more plain that deciment is unch combination being obvious to a particular whiled in the ert. "T" leter document published after the international filling data or priority date and not in conflict with the application but cated to understand the principle or lineary underlying the invention "4" document member of the same patent family Special categories of cited documents: 13
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"Or document referring to an oral disclosure, use, ashibition or other meens. "P" document published proof to the international filing dete but later than the priority date claimed

Date of Mailing of this International Search Report 9 Date of the Actual Completion of the International Search 1 IV, CERTIFICATION

18 December 1990 nternational Searching Authority ISA/US

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Richard Lebovitz

Form PCT/ISA/210 (second sheet) (May 1988)

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Attachment To PCT/IPEA/210 Classification Of Subject Matter

IPC(5): C12N 15/00; C12N 15/10; C07K 7/00; C07H 15/12 U.S. CL: 435/172.3; 536/27; 530/350

International Application No. PCT/US90/05682

4. The all seschable clams could be searched without effort justifying an additional lee, the international Searching Authority aid half more payment of the additional fee. 3. The required additional search least were timely gaid by the applicant. Consequently, this misernational search sport is seaffrored to the mineral the claims; it is covered by claim numbers: 1. O As at recursed additional search less were timely paid by the applicant, this international search report covers an searchable claims of the international septication. 2. As only some of the required sedillional search (sea were limitly paid by the applicant, this international kwa. : "sport control only those claims of the international spplication for which leas were paid, specifically claims: E Claim numbers . because they relate to parts of the international application that do nat compty with the preaccibed requirements to such an estent has no meaningful international search can be carried and 1, specifically: This international search report has not been essablished in respect of certain claims under Article 17(2) (a) for the following reasons: because they are dependent dains not drahad in accordance with the second and third seniences of . because they relate to subject matter I not required to be searched by this Authority, namely: 5-9,15 21,23 FURTHER INFORMATION CONTINUED FROM THE SECOND SHEFT

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No protest accompanied the payment of additional search fees. document. 3. Claim numbers_ PCT Rute 6.4(a). 1. Ctaim numbers Remark on Protest

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